

BEST AVAILABLE COPY

Helping Gail w/ subcloning

Looked at the cells.

- (c) Yesterday: transfections done made a marvelous recovery. They are nearly confluent & look great. The only exception is plates 2A & 2B. They show toxicity - probably due to the high lipofectin concentration. Time of lipofectin may also be an issue.
- (b) P34 & P26 cells look good - ready to pass Monday.

Gail appears to have converged in PAE1 sp1A. To combine

- grow up clone
- more restriction sites
- transfect LNCaP for p-gal activity.

Pass cells & set-up new plates

20 x 6 cm plates for tomorrow

9 x 150 cm boxes for freezing down

3 x 150 cm boxes for carrying

Gail is making minis and doing restriction digests. She

has 6 BstII clones for HindIII, 22 clones of 6 kb HindIII PstI in CN70 (p-gal in B5).

→ Made 10 ml of 10x ligation (without ATP) per protocol in computer - report of 10x 2 med of ligation & project. Sterile filtered. Froze ~~that~~ 1 ml aliquots

5.0 ml 1M Tris (7.6)

1.0 ml 1M MgCl₂

1.0 ml 0.5M DTT

50 µl 10 mg/ml BSA

2.95 ml H₂O

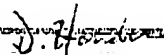
To Page No. _____

Witnessed & Understood by me,



Invented by

Recorded by



BEST AVAILABLE COPY

CNT10 - 1000 - 1000
CNT1 - 1000 - 1000

abund

PSC
580bp

AGENDA

- I. Characterization of PSE - Eric, Lena, Joe
5' end and 3' end of PSE defined within 200 bp
ARE within the Cla I site required
Sequencing
Thus, at least three elements: ARE, Left and Right
within PSE

Movement and orientation of minimal PSE - Lena (4 of 4)
Are these tested in LncAP yet?

Joe has sequenced my (CN42) Ck1 (Kramer) CN68

Gel Shift with 200 bp segments across PSE - Eric

fragment which spans the 5' end of the XbaI site. Linker protein from

- II. Transgenic Mice Constructs - Gall, Dan
Gall 1. 6kb driving β -gal (need 6 kb in CN70, β -gal in BS)
Lena 2. minimal enhancer driving β -gal
we have it 3. 6kb driving DT-A (CN45) Madrasa veta mouse

- III. Adenovirus Constructs - Gall, Lena, Dan

1. PSE minimal enhancer driving β -gal
2. PSE minimal enhancer driving CAT
3. PSE minimal enhancer driving DT-A

4. CMV driving HSV-tk done } for competent Y, if we add
5. CMV driving cytosine deaminase } drug we can kill virus
6. CMV driving β -gal positive control

Construction of Adenovirus vector

AE1sp1A AE1sp1B, BHG10, BHG11, BHG12, FG140, PXC1,
PABS.4

AE1A & BHG10
break virus within
Co-transfected & grow
recombined in cells.

- IV. Tissue Specificity of Minimal Enhancer - In Vitro - Eric and Joe
Results good result of last week being reported this week

- V. Tissue Specificity of PSE - In Vivo - Eric, Joe, Lena

Direct Injection of DNA into Tumors } exp. too big exp
Technique of Intra-tumor injection } exp ongoing
CAT Assay

- VI. Liposomes - Henry

Experiments

Commercial vs. Jensen low vs. Dan lower

Use commercial cells for next ligations

4/15 Cory Gorman from Megalabs talking about Liposomes
Henry in Tucson Mon. & Tues.

Check lunch schedule of Conf. room for virus lesson

check 3H allosteric - Delivering today! Friday lunch

Q for Eric IPA salt out. for removal of Eth Br. from CsCl/Eth Br./DNA from

Water solvents IPA - if water
is B. high salt & IPA is in salt
Water cannot solvate IPA
& you get an interface

To Page No.

Submitted by

Date

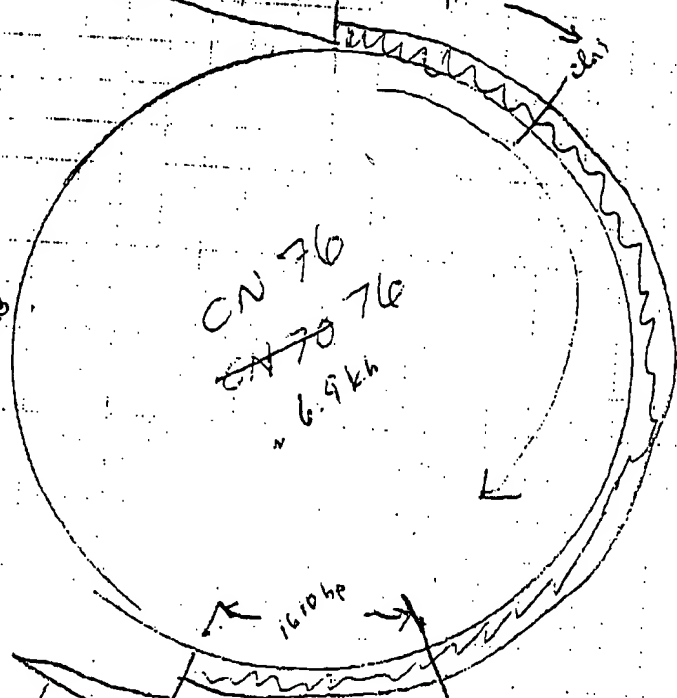
Recorded by

BEST AVAILABLE COPY

From Page No. 11

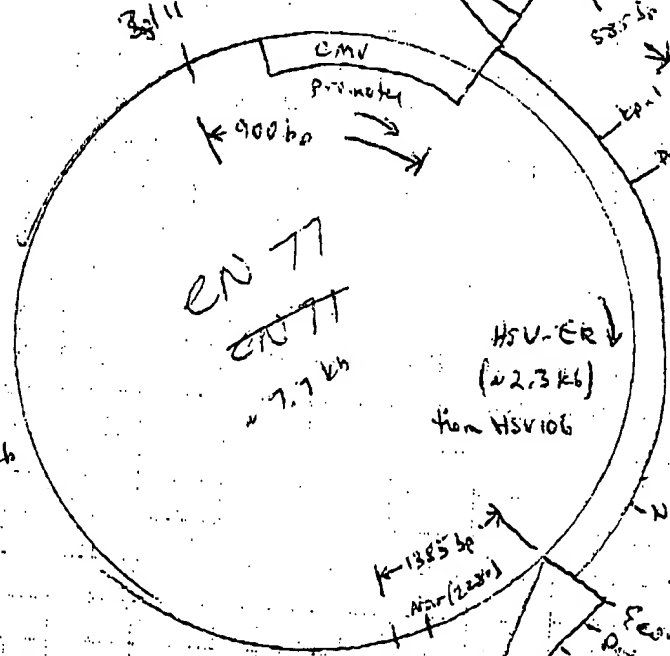
11
 12
 13
 14
 15
 16
 17
 18
 19
 20
 21
 22
 23
 24
 25
 26
 27
 28
 29
 30
 31
 32
 33
 34
 35
 36
 37
 38
 39
 40
 41
 42
 43
 44
 45
 46
 47
 48
 49
 50
 51
 52
 53
 54
 55
 56
 57
 58
 59
 60
 61
 62
 63
 64
 65
 66
 67
 68
 69
 70
 71
 72
 73
 74
 75
 76
 77
 78
 79
 80
 81
 82
 83
 84
 85
 86
 87
 88
 89
 90
 91
 92
 93
 94
 95
 96
 97
 98
 99
 100

35
 2.986 kb

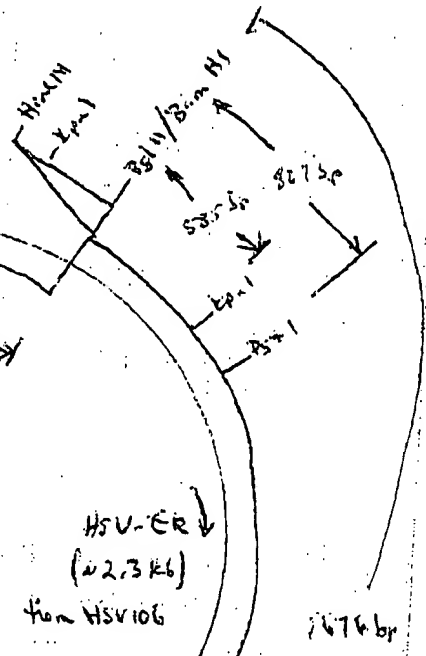


insert from pCMV A
 β-galactosidase
 = 3832 bp
 on XbaI fragment
 is promoterless

BamHI
 354X2 SmaI
 EcoRI NheI
 KpnI



pDNA3
 5.4 kb



number in () etc
 designation in pDNA3

cmv promoter driving
 HSV-ta in pDNA3

HSV-ER
 (2.3 kb)
 from HSV106
 1385 bp
 2333 bp
 919 bp

Witnessed & Understood by me,

[Signature]

Invented by

Recorded by

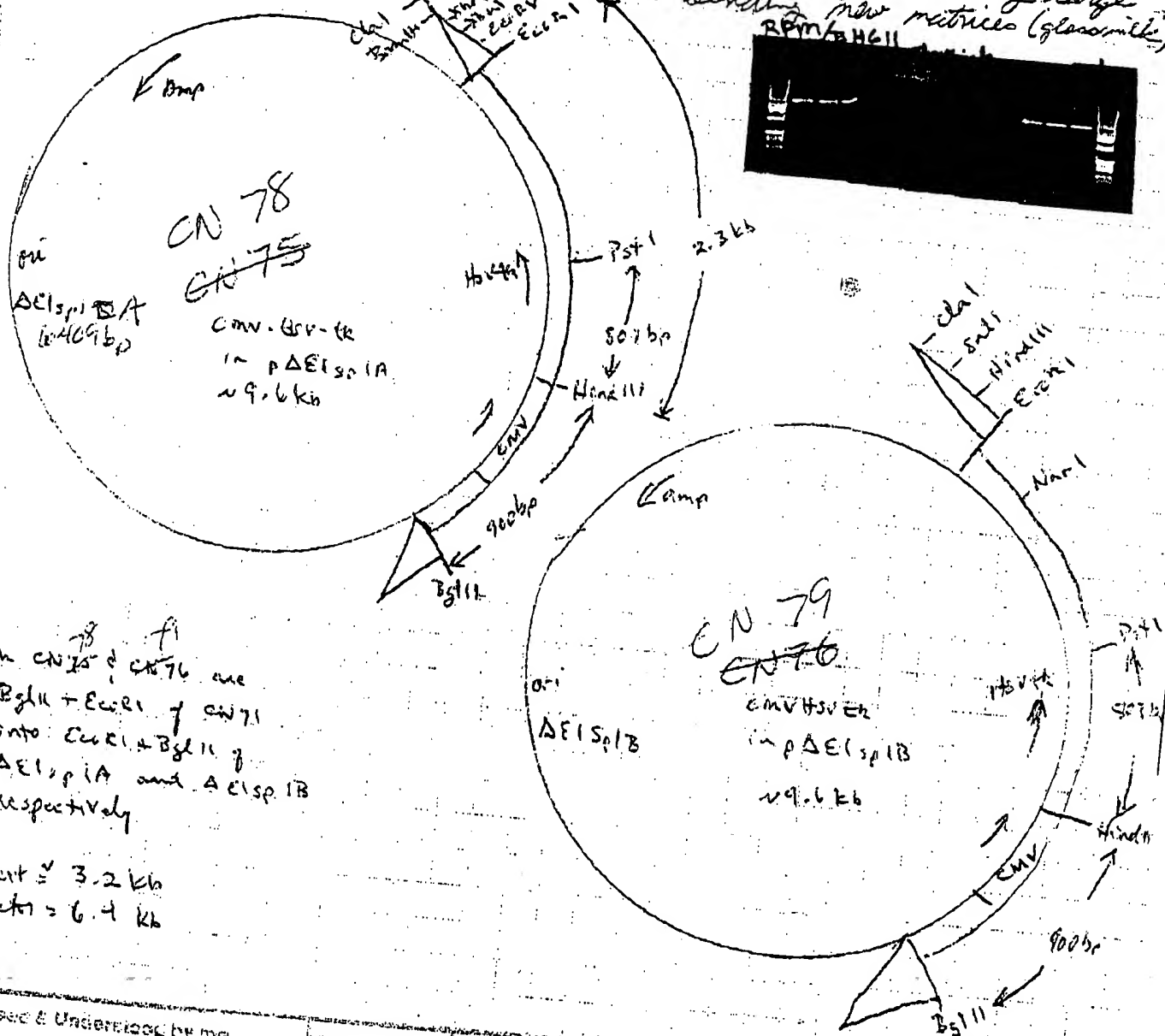
Date

To Page No.

Don has grown BHG-11 another A, B, C
cultures to HindIII screen.

I am now prepping 1.5 ml of
8 µl DNA
1 µl #2 10x Buffer
1 µl HindIII 2 µl - No luck

I called Bio 101 with regard to the problems I'm having with very large
plasmids in their RPM kit. They are sending new matrices (glass milk)
RPM/BHG11



Both CN 78 & CN 79 are
BglII + EcoRI of CN 71
into EcoRI + BglII of
ΔE1sp1A and ΔE1sp1B
respectively.

Insert = 3.2 kb
Vector = 6.4 kb

Witnessed & Understood by me.

[Signature]

Witnessed by

Witnessed by

To Page No.

Date

BEST AVAILABLE COPY

Book No. 16

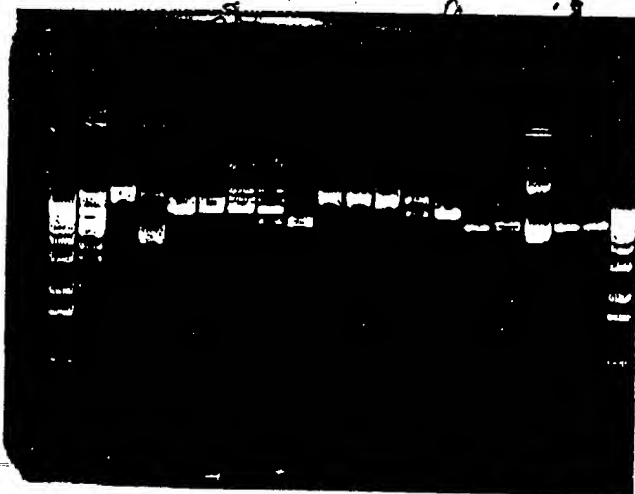
127

Concentrations

	OD260	OD280	260/280	Conc	Yield
FL140 16.122	0.076	0.036	2.1	380 µg/ml	152 µg
⁷⁸ CN 78-1 16.122	0.015	0.002	7.5	75 µg/ml	15 µg
⁷⁸ CN 78-2 16.122	0.437	0.208	2.1	220 µg/ml	437 µg
⁷⁹ CN 79-1 16.122	0.022	0.012	1.8	110 µg/ml	22 µg
⁷⁹ CN 79-2 16.122	0.264	0.133	2	1.3 mg/ml	264 µg

⁷⁸
CN 78 looks fine. 1X cut of HindIII, EcoRI, BglII, EcoRI+BglII given insert and HindIII cuts the insert into 2.3 kb if 900 bp as it should

⁷⁹
CN 79 may be fine but: it appears NruI and HindIII (lanes 10 & 11) were omitted: ie. no cut. But EcoRI cuts 1X, BglII cuts 1X, EcoRI+BglII give the 3.2 kb insert, and HindIII cuts the insert into 2-3 (visible in photo) 900 (visible by eye but not in photo) as it should



To Page No.

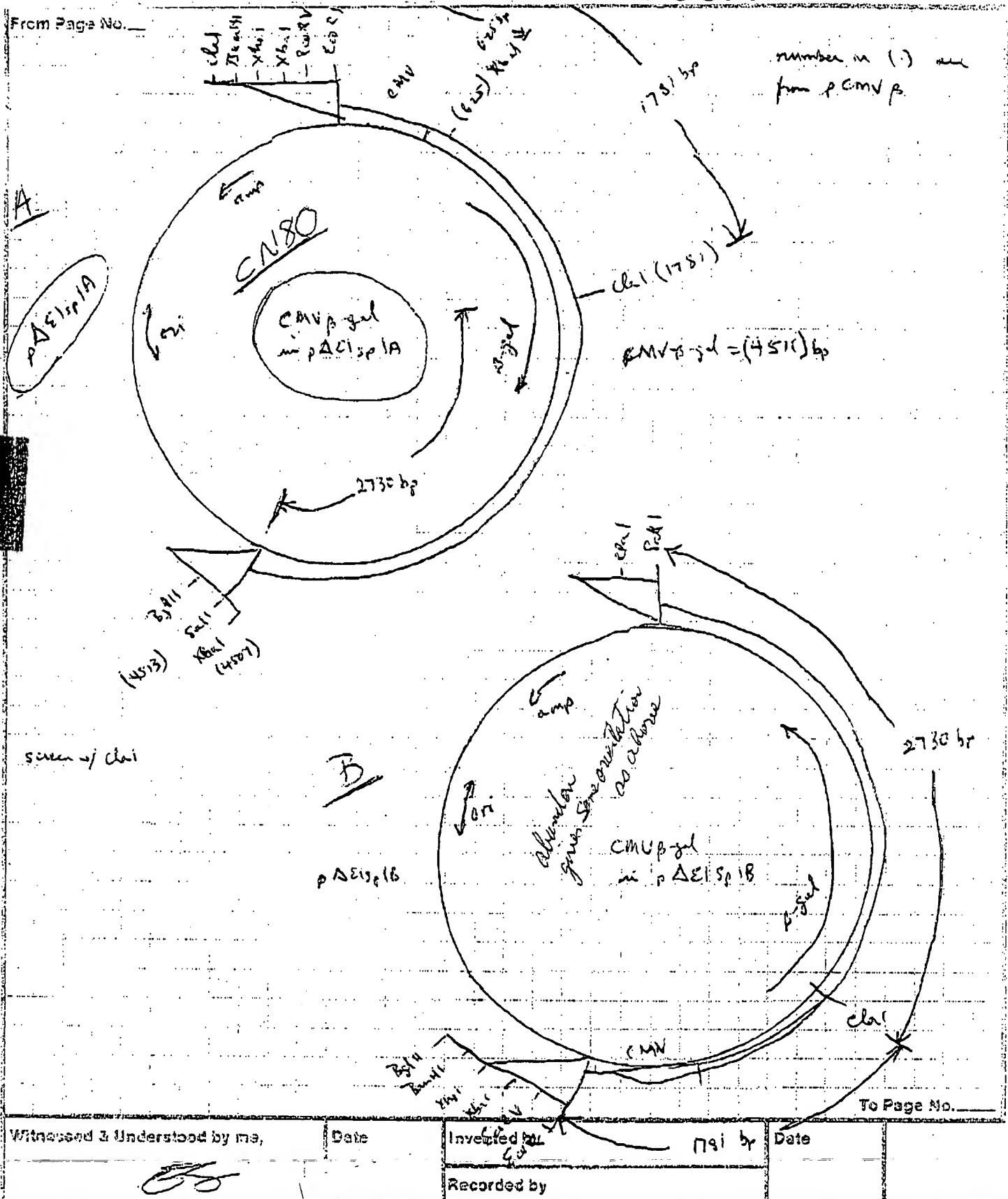
Date

Recorded by

E5

BEST AVAILABLE COPY

From Page No.



Witnessed & Understood by me,

Date _____

Invested for
G. J.

791 54

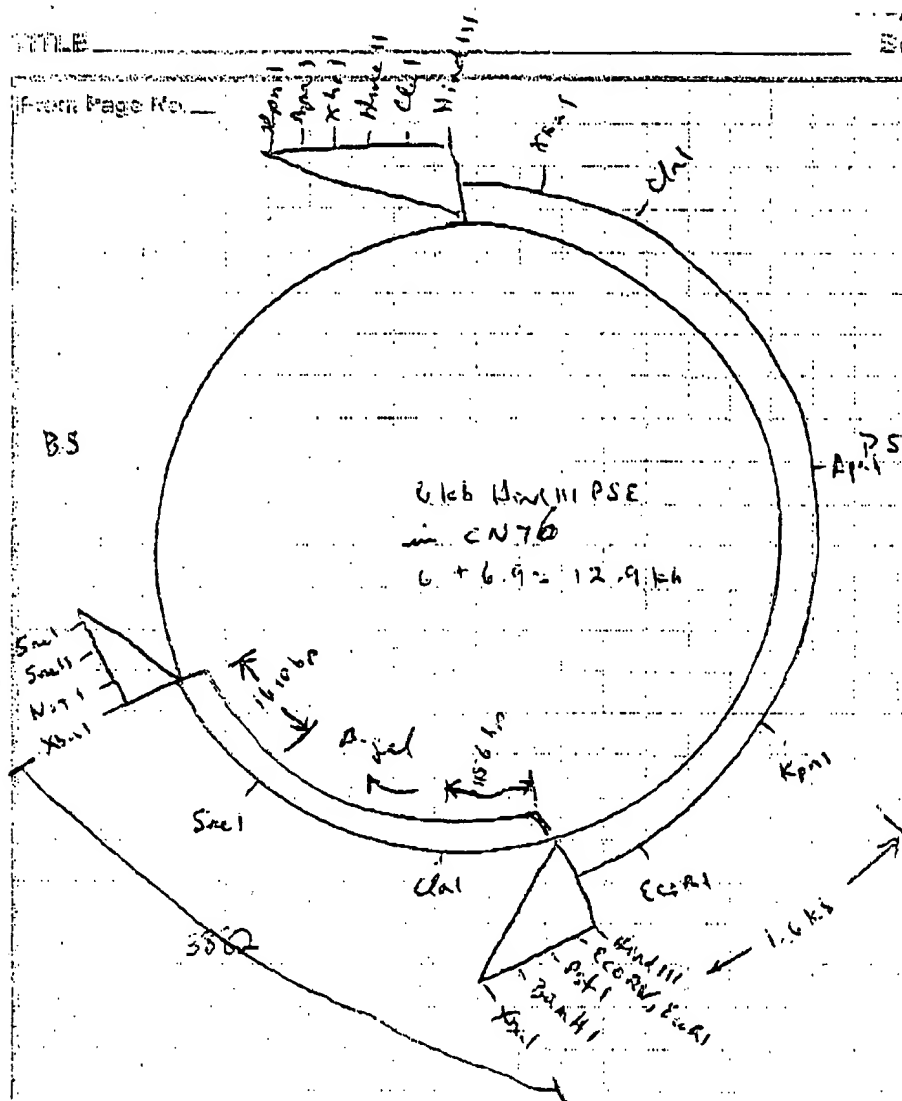
Date _____

To Page No.

Recorded by

777. 陸

Life: Page No. _____



Abel 1 7.55, 3.9, ~~2.4~~ 3.4
- 0.5, 3.9, ~~2.8~~ 8.4

$$\begin{array}{r} \text{Cl}_2 + 18,27,6 + 54,56 \\ - 3,4,2,5,6 \end{array}$$

April + 3.2, 9.7

Kpn.1 + 4,4,8.5

- 116, 113.

LA 75E (a 6 inch)

Am 811 - 660⁺ 6782.

Зачека w/krn1

if correct orientation
should get ~ 4.4 kb
if opposite orientation
should get ~ 1.6 kb

To Page No.

Witnessed & Understood by me.

Invented by

Deze

Recorded by

From 16.12.3

Don picked more 22 cultures on lig 16.12.3 this weekend. Of the 6 cultures I miniced the weekend I was correct but wrong quantities. There is a clone we will get something out of these 22.
Mini RPM + KpnI cut

4 μ l dH₂O
1 μ l 10xBSA
1 μ l 10xNEB1
1 μ l KpnI 20 μ l
3 μ l DNA

Mini RPM to TBamp cultures of BHK11 - 2 dif plates 1 commercial DNA 1 from previously good mini. These were picked on Sat 2 expanded to 10ml each. Don collected 1/2 ml for minis & fed ea culture 10ml TBamp.

16.12.3 prep
2 μ l

Clone gel Vectors + Inserts 16.12.3



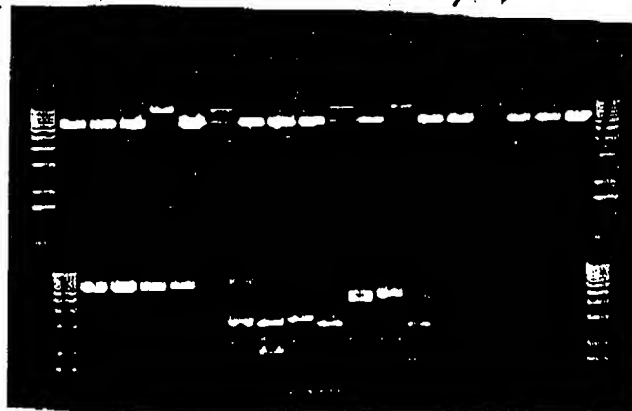
CATCH 4.111 prep A+B back equal 0 = 1/15 of 1 μ g ladder
Fr 1/15 = 0.066... 0.066/2 μ l = 33 ng/ μ l

Don did RE digest of SV CAT 16.78 see bank 15.123 this prep look good

OD 260 0.550
OD 280 0.321
260/280 1.7

Conc = 2.8 mg/ μ l

Don ~~not~~ digests of 16.12.3 above
close 3.5, 5 shows small band
at 4.9 but it is very light
I don't know whether to believe
it or not. I will do more
cuts tomorrow



To Page No.

Witnessed & Understood by me,

Date

Invented by

Date

Recorded by

[Signature]